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Bringing MMFF to the RDKit

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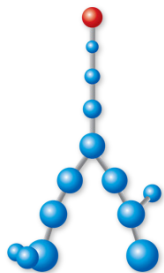
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(Article begins on next page)



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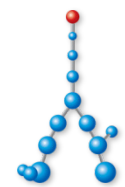


Bringing MMFF to the RDKit

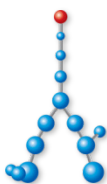
Paolo Tosco

RDKit User Group Meeting 2013

October 2–4, European Bioinformatics Institute, Hinxton, UK

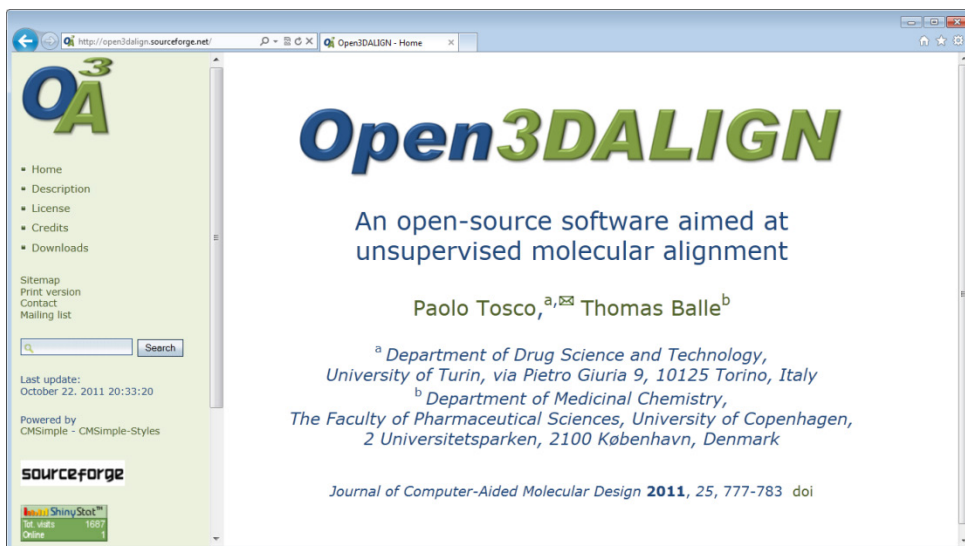


- The MMFF implementation
- UFF gains something, too
- MMFF applications



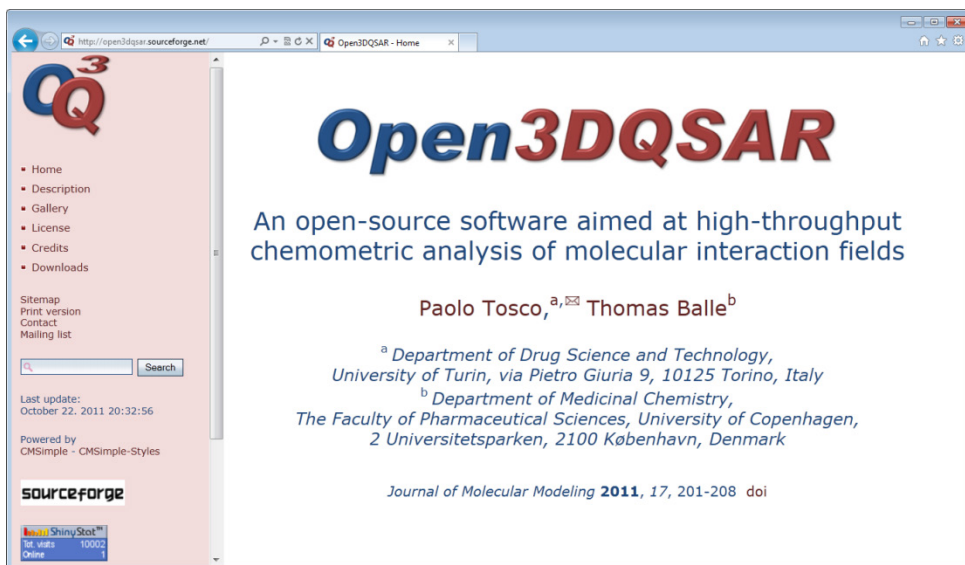
• Open3DALIGN

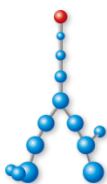
- conformational search
- molecular alignment
 - single/multiple conformations
 - atom-based/pharmacophore-based/mixed alignment
 - single-run/iterative



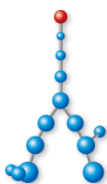
• Open3DQSAR

- MIF computation
- PLS model building and validation
- variable selection





- Preliminary step before implementing **Open3DQSAR** and **Open3DALIGN** functionality
- In principle, only MMFF atom types and charges are needed for that purpose...
- ...but one shouldn't be lazy, right? ;-)



Merck Molecular Force Field. I. Basis, Form, Scope, Parameterization, and Performance of MMFF94*

Merck Molecular Force Field. II. MMFF94 van der Waals and Electrostatic Parameters for Intermolecular Interactions*

Merck Molecular Force Field. III. Molecular Geometries and Vibrational Frequencies for MMFF94*

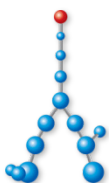
Merck Molecular Force Field. IV. Conformational Energies and Geometries for MMFF94*

Merck Molecular Force Field. V. Extension of MMFF94 Using Experimental Data, Additional Computational Data, and Empirical Rules*

MMFF VI. MMFF94s Option for Energy Minimization Studies

MMFF VII. Characterization of MMFF94, MMFF94s, and Other Widely Available Force Fields for Conformational Energies and for Intermolecular-Interaction Energies and Geometries

MMFF94(s) is fully documented by a series of **7** papers published by **Halgren** and Nachbar (Merck) on *J. Comp. Chem.* between 1995 and 1998

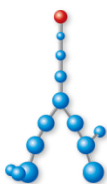


$$\begin{aligned}
 E_{\text{MMFF}} = & \sum \text{EB}_{ij} && \text{bond stretching} && \text{EB}_{ij} = 143.9325 \frac{kb_{ij}}{2} \Delta r_{ij}^2 \left(1 + cs \Delta r_{ij} + \frac{7}{12} cs^2 \Delta r_{ij}^2 \right) \\
 & + \sum \text{EA}_{ijk} && \text{angle bending} && \text{EA}_{ijk} = 0.043844 \frac{ka_{ijk}}{2} \Delta \vartheta_{ijk}^2 \left(1 + cb \Delta \vartheta_{ijk} \right) \\
 & + \sum \text{EBA}_{ijk} && \text{stretch-bend} && \text{EBA}_{ijk} = 2.51210 (kba_{ijk} \Delta r_{ij} + kba_{kjl} \Delta r_{kj}) \Delta \vartheta_{ijk} \\
 & + \sum \text{EOOP}_{ijk;l} && \text{out-of-plane bending} && \text{EOOP}_{ijk;l} = 0.043844 \frac{k\text{oop}_{ijk;l}}{2} \chi_{ijk;l}^2 \\
 & + \sum \text{ET}_{ijkl} && \text{torsion} && \text{ET}_{ijkl} = 0.5 [V_1 (1 + \cos \phi) + V_2 (1 - \cos 2\phi) + V_3 (1 + \cos 3\phi)] \\
 & + \sum \text{EvdW}_{ij} && \text{van der Waals} && \text{EvdW}_{ij} = \varepsilon_{ij} \left(\frac{1.07 R_{ij}^*}{R_{ij} + 0.07 R_{ij}^*} \right)^7 \left(\frac{1.12 R_{ij}^{*7}}{R_{ij}^7 + 0.12 R_{ij}^{*7}} - 2 \right) \\
 & + \sum \text{EQ}_{ij} && \text{electrostatic} && \text{EQ}_{ij} = 332.0716 \frac{q_i q_j}{D (R_{ij} + \delta)^n}
 \end{aligned}$$

CR	1	ALKYL CARBON, SP3	HO	21	GENERAL H ON OXYGEN	NPO2	43	NITROGEN IN PHOSPHONAMIDES
C=C	2	VINYLIC CARBON, SP2	HOM	21	HYDROGEN IN HYDROXIDE ANION	NPO3	43	NITROGEN IN PHOSPHONAMIDES, THREE O'S ON P
CSP2	2	GENERIC SP2 CARBON	CR3R	22	CARBON IN A 3-MEMBERED RING	NC%N	43	NITROGEN ATTACHED TO CYANO GROUP
C=O	3	GENERAL CARBONYL CARBON	HNR	23	H-N (SP3)	STHI	44	SULFUR AS IN THIOPHENE
C=N	3	SP2 CARBON IN C=N	H3N	23	H-N (SP3), AMMONIA	NO2	45	NITRO GROUP NITROGEN
CGD	3	GUANIDINE CARBON, DOUBLY BONDED TO N	HPYL	23	H-N IN PYRROLE	NO3	45	NITRATE GROUP NITROGEN
C=OR	3	KETONE OR ALDEHYDE CARBONYL CARBON	HNOX	23	H-N IN IN A N-OXIDE	N=O	46	NITROSO NITROGEN
C=ON	3	AMIDE CARBONYL CARBON	HNM	23	H ON DICOORD, NEGATIVELY CHARGED NITROGEN	NAZT	47	TERMINAL NITROGEN IN AZIDO OR DIAZO GROUP
CONN	3	UREA CARBONYL CARBON	HN	23	GENERAL H ON NITROGEN	NSO	48	DIVALENT NITROGEN REPLACING MONOVALENT O IN SO2 GROUP
COO	3	CARBOXYLIC ACID OR ESTER CARBONYL CARBON	HOCO	24	H-O IN CARBOXYLIC ACIDS	O+	49	POSITIVELY CHARGED OXONIUM (TRICOORDINATE) OXYGEN
COON	3	CARBAMATE CARBONYL CARBON	HOP	24	HYDROGEN ON OXYGEN ATTACHED TO PHOSPHOROUS	HO+	50	HYDROGEN ON O+ OXYGEN
COOO	3	C ARBONIC ACID OR ESTER CARBONYL CARBON	PO4	25	PHOSPHOROUS IN PHOSPHATES AND PHOSPHODIESTERS	O=+	51	POSITIVELY CHARGED OXENIUM (DICOORDINATE) OXYGEN
C=OS	3	THIOESTER CARBONYL CARBON, DOUBLE BONDED TO O	PO3	25	TETRACOORDINATE P WITH THREE ATTACHED OXYGENS	HO=+	52	HYDROGEN ON OXENIUM OXYGEN
C=S	3	THIOESTER CARBON, DOUBLY BONDED TO S	PO2	25	TETRACOORDINATE P WITH TWO ATTACHED OXYGENS	=N=	53	NITROGEN IN C=N=N OR -N=N=N
C=SN	3	THIOAMIDE, CARBON, DOUBLY BONDED TO S	PO	25	TETRACOORDINATE P WITH ONE ATTACHED OXYGEN	N+=C	54	POSITIVELY CHARGED IMINIUM NITROGEN
CSO2	3	CARBON IN >C=SO2	PTET	25	GENERAL TETRACOORDINATE PHOSPHORUS	N+=N	54	POSITIVELY CHARGED NITROGEN DOUBLE-BONDED TO N
CS=O	3	CARBON IN >C=S=O (SULFINYL GROUP)	P	26	TRICOORDINATE P, AS IN PHOSPHINES	NCN+	55	N IN +N=C-N RESONANCE STRUCTURES - FORMAL CHARGE=1/2
CSS	3	THIOCARBOXYLIC ACID OR ESTER CARBONYL CARBON	HN=N	27	AZO HYDROGEN	NGD+	56	GUANIDINIUM-TYPE NITROGEN - FORMAL CHARGE=1/3
C=P	3	CARBON DOUBLE BONDED TO PHOSPHOROUS	HN=C	27	IMINE HYDROGEN	CGD+	57	GUANIDINIUM CARBON
CSP	4	ACETYLENIC CARBON	HNCO	28	AMIDE HYDROGEN	CNN+	57	C IN +N=C-N RESONANCE STRUCTURES
=C=	4	ALLENIC CARBON	HNCS	28	THIOAMIDE HYDROGEN	NPd+	58	PYRIDINIUM-TYPE NITROGEN - FORMAL CHARGE=1
HC	5	H ATTACHED TO C	HNCC	28	H-N IN ENAMINES	OFUR	59	AROMATIC OXYGEN AS IN FURAN
HSI	5	H ATTACHED TO SI	HNCN	28	H-N IN H-N-C=N	C%	60	ISONITRILE CARBON
OR	6	ALCOHOL OR ETHER OXYGEN	HNNC	28	H-N IN H-N-N=C	NR%	61	ISONITRILE NITROGEN [FC = 0] OR DIAZO NITROGEN [FC=1]
OC=O	6	ESTER OR CARBOXYLIC ACID -O-	HNNN	28	H-N IN H-N-N=N	NM	62	DEPROTONATED SULFONAMIDE N-; FORMAL CHARGE=-1
OC=C	6	ENOLIC OR PHENOLIC OXYGEN	HNSO	28	H-N IN SULFONAMIDE	C5A	63	ALPHA CARBON IN 5-MEMBERED HETEROAROMATIC RING
OC=N	6	DIVALENT OXYGEN	HNPO	28	H-N IN PHOSPHONAMIDE	C5B	64	BETA CARBON IN 5-MEMBERED HETEROAROMATIC RING
OC=S	6	THIOESTER OR THIOACID -O-	HNC%	28	HYDROGEN ON N ATTACHED TO TRIPLY BONDED CARBON	N5A	65	ALPHA AROM HETEROCYCLIC 5-RING NITROGEN
ONO2	6	DIVALENT NITRATE "ETHER" OXYGEN	HSP2	28	GENERAL H ON SP2 NITROGEN	N5B	66	BETA AROM HETEROCYCLIC 5-RING NITROGEN
ON=O	6	DIVALENT NITRITE "ETHER" OXYGEN	HOCc	29	H-O IN ENOLS AND PHENOLS	N2OX	67	SP2-HYDRIDIZED N-OXIDE NITROGEN
OSO3	6	DIVALENT OXYGEN ATTACHED TO SULFUR	HOCN	29	H-O IN HO-C=N	N3OX	68	SP3-HYDRIDIZED N-OXIDE NITROGEN
OSO2	6	DIVALENT OXYGEN ATTACHED TO SULFUR	CE4R	30	OLEFINIC CARBON IN 4-MEMBERED RINGS	NPOX	69	PYRIDINE N-OXIDE NITROGEN
OSO	6	DIVALENT OXYGEN ATTACHED TO SULFUR	HOH	31	HYDROGEN IN H2O	OH2	70	OXYGEN ON WATER
OS=O	6	DIVALENT OXYGEN ATTACHED TO SULFOXIDE SULFUR	O2CM	32	OXYGEN IN CARBOXYLATE ANION	HS	71	H ATTACHED TO DIVALENT, DICOORDINATE S
-OS	6	GENERAL DIVALENT OX ATTACHED TO S	OXN	32	N-OXIDE OXYGEN	HS=N	71	H ATTACHED TO TETRAVALENT, TRICOODR S DBL BONDED TO N
OPO3	6	DIVALENT OXYGEN ATTACHED TO PHOSPHOROUS	O2N	32	NITRO OXYGEN	HP	71	H ATTACHED TO TRI- OR TETRACOORDINATE PHOSPHORUS
OPO2	6	DIVALENT OXYGEN ATTACHED TO PHOSPHOROUS	O2NO	32	NITRO-GROUP OXYGEN IN NITRATE	S-P	72	TERMINAL SULFUR BONDED TO PHOSPHORUS
OPO	6	DIVALENT OXYGEN ATTACHED TO PHOSPHOROUS	O3N	32	NITRATE ANION OXYGEN	S2CM	72	TERMINAL SULFUR IN THIOCARBOXYLATE ANION
-OP	6	DIVALENT OXYGEN ATTACHED TO PHOSPHOROUS	O-S	32	SINGLE TERMINAL OXYGEN ON TETRACOORD SULFUR	SM	72	TERMINAL SULFUR - FORMAL CHARGE=-1
-O-	6	GENERAL DIVALENT O	O2S	32	TERMINAL O-S IN SULFONES AND SULFONAMIDES	SSMO	72	TERMINAL SULFUR IN THIOSULFATE GROUP
O=C	7	GENERAL C=O	O3S	32	TERMINAL O IN SULFONATES	SO2M	73	SULFUR IN NEGATIVELY CHARGED SULFINATE GROUP
O=CN	7	CARBONYL OXYGEN, AMIDES	O4S	32	TERMINAL O IN SO4(-3)	SSOM	73	TRICOORD SULFUR IN THIOSULFINATE GROUP
O=CR	7	CARBONYL OXYGEN, ALDEHYDES AND KETONES	OSMS	32	TERM O IN THIOSULFINATE ANION - FORMAL CHARGE=-0.5	=S=O	74	SULFINYL SULFUR, EG. IN C=S=O
O=CO	7	CARBONYL OXYGEN, CARBOXYLIC ACIDS AND ESTERS	OP	32	TERMINAL O IN PHOSPHOXIDES	-P=C	75	PHOSPHOROUS DOUBLY BONDED TO CARBON
O=N	7	NITROSO OXYGEN	O2P	32	TERMINAL O IN PHOSPHINATES	N5M	76	NEGATIVELY CHARGED N IN, E.G., TRI- OR TETRAZOLE ANION
O=S	7	O=S IN SULFOXIDES	O3P	32	TERMINAL OXYGEN IN PHOSPHONATES	CLO4	77	CHLORINE IN PERCHLORATE ANION, CLO4(-)
O=S=	7	O=S ON SULFUR DOUBLY BONDED TO, E.G., CARBON	O4P	32	TERMINAL OXYGEN IN PHOSPHATES AND PHOSPHODIESTERS	C5	78	GENERAL CARBON IN 5-MEMBERED HETEROAROMATIC RING
NR	8	NITROGEN IN ALIPHATIC AMINES	O4CL	32	OXYGEN IN CLO4(-) ANION - FORMAL CHARGE=-0.25	N5	79	GENERAL NITROGEN IN 5-MEMBERED HETEROCYCLIC RING
N=C	9	NITROGEN IN IMINES	HOS	33	H ON OXYGEN ATTACHED TO SULFUR	CIM+	80	C IN N-C-N IN IMIDAZOLIUM ION
N=N	9	NITROGEN IN AZO COMPOUNDS	NR+	34	QUATERNARY NITROGEN, SP3, POSITIVELY CHARGED	NIM+	81	IMIDAZOLIUM-TYPE NITROGEN - FORMAL CHARGE=1/2
NC=O	10	NITROGEN IN AMIDES	OM	35	ALKOXIDE OXYGEN, NEGATIVELY CHARGED	N5A+	81	POSITIVE N5A NITROGEN - FORMAL CHARGE=1
NC=S	10	NITROGEN IN N-C=S, THIOAMIDE	OM2	35	OXIDE OXYGEN ON SP2 CARBON, NEGATIVELY CHARGED	N5B+	81	POSITIVE N5B NITROGEN - FORMAL CHARGE=1
NN=C	10	NITROGEN IN N=N=C	HNR+	36	H ON QUATERNARY NITROGEN	N5+	81	POSITIVE N5 NITROGEN - FORMAL CHARGE=1
NN=N	10	NITROGEN IN N=N=N	HIM+	36	H ON IMIDAZOLIUM-TYPE NITROGEN	N5AX	82	N-OXIDE NITROGEN IN 5-RING ALPHA POSITION
F	11	FLUORINE	HPD+	36	H ON PROTONATED PYRIDINE NITROGEN	N5BX	82	N-OXIDE NITROGEN IN 5-RING BETA POSITION
CL	12	CHLORINE	HNN+	36	H ON AMIDINIUM-TYPE NITROGEN	N5OX	82	N-OXIDE NITROGEN IN GENERAL 5-RING POSITION
BR	13	BROMINE	HNC+	36	H ON PROTONATED IMINE NITROGEN	FE+2	87	IRON +2 CATION
I	14	IODINE	HGD+	36	H ON GUANIDINIUM-TYPE NITROGEN	FE+3	88	IROM +3 CATION
S	15	SULFUR IN THIOETHERS AND MERCAPTANS	HN5+	36	H ON N5+, N5A+ OR N5B+	F-	89	FLUORIDE ANION
S=C	16	TERMINAL SULFUR DOUBLY BONDED TO CARBON	CB	37	CARBON AS IN BENZENE, PYRROLE	CL-	90	CHLORIDE ANION
S=O	17	SULFUR IN SULFOXIDES	NPYD	38	NITROGEN, AS IN PYRIDINE	BR-	91	BROMIDE ANION
>S=N	17	SULFUR, TRICOORD, DOUBLY BONDED TO N	NPYL	39	NITROGEN, AS IN PYRROLE	LI+	92	LITHIUM CATION
SO2	18	SULFUR IN SULFONES	NC=C	40	NITROGEN ON N-C=C	NA+	93	SODIUM CATION
SO2N	18	SULFUR IN SULFONAMIDES	NC=N	40	NITROGEN IN N-C=N	K+	94	POTASSIUM CATION
SO3	18	SULFONATE SULFUR	NC=P	40	NITROGEN IN N-C=P	ZINC	95	DIPOSITIVE ZINC
SO4	18	SULFATE SULFUR	NC%C	40	NITROGEN ATTACHED TO C-C TRIPLE BOND	ZN+2	95	DIPOSITIVE ZINC
=SO2	18	SULFONE SULPHER DOUBLY BONDED TO CARBON	CO2M	41	CARBOXYLATE ANION CARBON	CA+2	96	DIPOSITIVE CALCIUM
SNO	18	SULFUR IN NITROGEN ANALOG OF A SULFONE	CS2M	41	CARBON IN THIOCARBOXYLATE ANION	CU+1	97	MONOPOSITIVE COPPER
SI	19	SILICON	NSP	42	NITROGEN, TRIPLE BONDED	CU+2	98	DIPOSITIVE COPPER
CR4R	20	CARBON IN 4-MEMBERED RINGS	NSO2	43	NITROGEN IN SULFONAMIDES	MG+2	99	DIPOSITIVE MAGNESIUM CATION
HOR	21	HYDROGEN IN ALCOHOLS	NSO3	43	NITROGEN IN SULFONAMIDES, THREE O'S ON S			

CR	1	ALKYL CARBON, SP3	HO	21	GENERAL H ON OXYGEN	NPO2	43	NITROGEN IN PHOSPHONAMIDES
C=C	2	VINYLIC CARBON, SP2	HOM	21	HYDROGEN IN HYDROXIDE ANION	NPO3	43	NITROGEN IN PHOSPHONAMIDES, THREE O'S ON P
CSP2	2	GENERIC SP2 CARBON	CR3R	22	CARBON IN A 3-MEMBERED RING	NC%N	43	NITROGEN ATTACHED TO CYANO GROUP
C=O	3	GENERAL CARBONYL CARBON	HNR	23	H-N (SP3)	STHI	44	SULFUR AS IN THIOPHENE
C=N	3	SP2 CARBON IN C=N	H3N	23	H-N (SP3), AMMONIA	NO2	45	NITRO GROUP NITROGEN
CGD	3	GUANIDINE CARBON, DOUBLY BONDED TO N	HPYL	23	H-N IN PYRROLE	NO3	45	NITRATE GROUP NITROGEN
C=OR	3	KETONE OR ALDEHYDE CARBONYL CARBON	HNOX	23	H-N IN IN A N-OXIDE	N=O	46	NITROSO NITROGEN
C=ON	3	AMIDE CARBONYL CARBON	HNM	23	H ON DICOORD, NEGATIVELY CHARGED NITROGEN	NAZT	47	TERMINAL NITROGEN IN AZIDO OR DIAZO GROUP
CONN	3	UREA CARBONYL CARBON	HN	23	GENERAL H ON NITROGEN	NSO	48	DIVALENT NITROGEN REPLACING MONOVALENT O IN SO2 GROUP
COO	3	CARBOXYLIC ACID OR ESTER CARBONYL CARBON	HOCO	24	H-O IN CARBOXYLIC ACIDS	O+	49	POSITIVELY CHARGED OXONIUM (TRICOORDINATE) OXYGEN
COON	3	CARBAMATE CARBONYL CARBON	HOP	24	HYDROGEN ON OXYGEN ATTACHED TO PHOSPHOROUS	HO+	50	HYDROGEN ON O+ OXYGEN
COOO	3	C ARBONIC ACID OR ESTER CARBONYL CARBON	PO4	25	PHOSPHOROUS IN PHOSPHATES AND PHOSPHODIESTERS	O=+	51	POSITIVELY CHARGED OXENIUM (DICOORDINATE) OXYGEN
C=OS	3	THIOESTER CARBONYL CARBON, DOUBLE BONDED TO O	PO3	25	TETRACOORDINATE P WITH THREE ATTACHED OXYGENS	HO=+	52	HYDROGEN ON OXENIUM OXYGEN
C=S	3	THIOESTER CARBON, DOUBLY BONDED TO S	PO2	25	TETRACOORDINATE P WITH TWO ATTACHED OXYGENS	=N=	53	NITROGEN IN C=N=N OR -N=N=N
C=SN	3	THIOAMIDE, CARBON, DOUBLY BONDED TO S	PO	25	TETRACOORDINATE P WITH ONE ATTACHED OXYGEN	N+=C	54	POSITIVELY CHARGED IMINIUM NITROGEN
CSO2	3	CARBON IN >C=SO2	PTET	25	GENERAL TETRACOORDINATE PHOSPHORUS	N+=N	54	POSITIVELY CHARGED NITROGEN DOUBLE-BONDED TO N
CS=O	3	CARBON IN >C=S=O (SULFINYL GROUP)	P	26	TRICOORDINATE P, AS IN PHOSPHINES	NCN+	55	N IN +N=C-N RESONANCE STRUCTURES - FORMAL CHARGE=1/2
CSS	3	THIOCARBOXYLIC ACID OR ESTER CARBONYL CARBON	HN=N	27	AZO HYDROGEN	NGD+	56	GUANIDIUM-TYPE NITROGEN - FORMAL CHARGE=1/3
C=P	3	CARBON DOUBLE BONDED TO PHOSPHOROUS	HN=C	27	IMINE HYDROGEN	CGD+	57	GUANIDIUM CARBON
CSP	4	ACETYLENIC CARBON	HNCO	28	AMIDE HYDROGEN	CNN+	57	C IN +N=C-N RESONANCE STRUCTURES
=C=	4	ALLENIC CARBON	HNCS	28	THIOAMIDE HYDROGEN	NPD+	58	PYRIDINIUM-TYPE NITROGEN - FORMAL CHARGE=1
HC	5	H ATTACHED TO C	HNCC	28	H-N IN ENAMINES	OFUR	59	AROMATIC OXYGEN AS IN FURAN
HSI	5	H ATTACHED TO SI	HNCN	28	H-N IN H-N-C=N	C%	60	ISONITRILE CARBON
OR	6	ALCOHOL OR ETHER OXYGEN	HNNC	28	H-N IN H-N-N=C	NR%	61	ISONITRILE NITROGEN [FC = 0] OR DIAZO NITROGEN [FC=1]
OC=O	6	ESTER OR CARBOXYLIC ACID -O-	HNNN	28	H-N IN H-N-N=N	NM	62	DEPROTONATED SULFONAMIDE N-; FORMAL CHARGE=-1
OC=C	6	ENOLIC OR PHENOLIC OXYGEN	HNSO	28	H-N IN SULFONAMIDE	C5A	63	ALPHA CARBON IN 5-MEMBERED HETEROAROMATIC RING
OC=N	6	DIVALENT OXYGEN	HNPO	28	H-N IN PHOSPHONAMIDE	C5B	64	BETA CARBON IN 5-MEMBERED HETEROAROMATIC RING
OC=S	6	THIOESTER OR THIOACID -O-	HNC%	28	HYDROGEN ON N ATTACHED TO TRIPLY BONDED CARBON	N5A	65	ALPHA AROM HETEROCYCLIC 5-RING NITROGEN
ONO2	6	DIVALENT NITRATE "ETHER" OXYGEN	HSP2	28	GENERAL H ON SP2 NITROGEN	N5B	66	BETA AROM HETEROCYCLIC 5-RING NITROGEN
ON=O	6	DIVALENT NITRITE "ETHER" OXYGEN	HOCC	29	H-O IN ENOLS AND PHENOLS	N2OX	67	SP2-HYDRIDIZED N-OXIDE NITROGEN
OSO3	6	DIVALENT OXYGEN ATTACHED TO SULFUR	HOCN	29	H-O IN HO-C=N	N3OX	68	SP3-HYDRIDIZED N-OXIDE NITROGEN
OSO2	6	DIVALENT OXYGEN ATTACHED TO SULFUR	CE4R	30	OLEFINIC CARBON IN 4-MEMBERED RINGS	NPOX	69	PYRIDINE N-OXIDE NITROGEN
OSO	6	DIVALENT OXYGEN ATTACHED TO SULFUR	HOR	31	HYDROGEN IN H2O	OH2	70	OXYGEN ON WATER
OS=O	6	DIVALENT OXYGEN ATTACHED TO SULFOXIDE SULFUR	O2M	32	TERMINAL OXYGEN ON TETRACORDINATE	OSR	71	H ATTACHED TO DIVALENT, DICOORDINATE S
-OS	6	GENERAL DIVALENT OX ATTACHED TO S	O2M	32	TERMINAL OXYGEN ON TETRACORDINATE	OSR	71	H ATTACHED TO TETRAVALENT, TRICOODR S DBL BONDED TO N
OPO3	6	DIVALENT OXYGEN ATTACHED TO PHOSPHOROUS	O2N	32	NITRO OXYGEN	P	71	H ATTACHED TO TRI- OR TETRACOORDINATE PHOSPHORUS
OPO2	6	DIVALENT OXYGEN ATTACHED TO PHOSPHOROUS	O2NO	32	NITRO-GROUP OXYGEN IN NITRATE	S-P	72	TERMINAL SULFUR BONDED TO PHOSPHORUS
OPO	6	DIVALENT OXYGEN ATTACHED TO PHOSPHOROUS	O3M	32	TERMINAL OXYGEN ON TETRACORDINATE	S-P	72	TERMINAL SULFUR IN THIOCARBOXYLATE ANION
-OP	6	DIVALENT OXYGEN ATTACHED TO PHOSPHOROUS	O2S	32	TERMINAL O-S IN SULFONES AND SULFONAMIDES	SSMO	72	TERMINAL SULFUR - FORMAL CHARGE=-1
-O-	6	GENERAL DIVALENT O	O3S	32	TERMINAL O IN SULFONATES	SO2M	73	SULFUR IN NEGATIVELY CHARGED SULFINATE GROUP
O=C	7	GENERAL C=O	O4S	32	TERMINAL O IN SO4(-3)	SSOM	73	TRICOORD SULFUR IN THIOSULFINATE GROUP
O=CN	7	CARBONYL OXYGEN, AMIDES	OSMS	32	TERM O IN THIOSULFINATE ANION - FORMAL CHARGE=-0.5	=S=O	74	SULFINYL SULFUR, EG. IN C=S=O
O=CR	7	CARBONYL OXYGEN, ALDEHYDES AND KETONES	OP	32	TERMINAL O IN PHOSPHOXIDES	-P=C	75	PHOSPHOROUS DOUBLY BONDED TO CARBON
O=CO	7	CARBONYL OXYGEN, CARBOXYLIC ACIDS AND ESTERS	O2P	32	TERMINAL O IN PHOSPHINATES	N5M	76	NEGATIVELY CHARGED N IN, E.G., TRI- OR TETRAZOLE ANION
O=N	7	NITROSO OXYGEN	O3P	32	TERMINAL OXYGEN IN PHOSPHONATES	CLO4	77	CHLORINE IN PERCHLORATE ANION, CLO4(-)
O=S	7	O=S IN SULFOXIDES	O4P	32	TERMINAL OXYGEN IN PHOSPHATES AND PHOSPHODIESTERS	C5	78	GENERAL CARBON IN 5-MEMBERED HETEROAROMATIC RING
O=S=	7	O=S ON SULFUR DOUBLY BONDED TO, E.G., CARBON	O4CL	32	OXYGEN IN CLO4(-) ANION - FORMAL CHARGE=-0.25	N5	79	GENERAL NITROGEN IN 5-MEMBERED HETEROCYCLIC RING
NR	8	NITROGEN IN ALIPHATIC AMINES	HOS	33	H ON OXYGEN ATTACHED TO SULFUR	CIM+	80	C IN N-C-N IN IMIDAZOLIUM ION
N=C	9	NITROGEN IN IMINES	NR+	34	QUATERNARY NITROGEN, SP3, POSITIVELY CHARGED	NIM+	81	IMIDAZOLIUM-TYPE NITROGEN - FORMAL CHARGE=1/2
N=N	9	NITROGEN IN AZO COMPOUNDS	OM	35	ALKOXIDE OXYGEN, NEGATIVELY CHARGED	N5A+	81	POSITIVE N5A NITROGEN - FORMAL CHARGE=1
NC=O	10	NITROGEN IN AMIDES	OM2	35	OXIDE OXYGEN ON SP2 CARBON, NEGATIVELY CHARGED	N5B+	81	POSITIVE N5B NITROGEN - FORMAL CHARGE=1
NC=S	10	NITROGEN IN N-C=S, THIOAMIDE	HNR+	36	H ON QUATERNARY NITROGEN	N5+	81	POSITIVE N5 NITROGEN - FORMAL CHARGE=1
NN=C	10	NITROGEN IN N=N=C	HIM+	36	H ON IMIDAZOLIUM-TYPE NITROGEN	N5AX	82	N-OXIDE NITROGEN IN 5-RING ALPHA POSITION
NN=N	10	NITROGEN IN N=N=N	HPD+	36	H ON PROTONATED PYRIDINE NITROGEN	N5BX	82	N-OXIDE NITROGEN IN 5-RING BETA POSITION
F	11	FLUORINE	HNN+	36	H ON AMIDINIUM-TYPE NITROGEN	N5OX	82	N-OXIDE NITROGEN IN GENERAL 5-RING POSITION
CL	12	CHLORINE	HNC+	36	H ON PROTONATED IMINE NITROGEN	FE+2	87	IRON +2 CATION
BR	13	BROMINE	HGD+	36	H ON GUANIDIUM-TYPE NITROGEN	FE+3	88	IROM +3 CATION
I	14	IODINE	HN5+	36	H ON N5+, N5A+ OR N5B+	F-	89	FLUORIDE ANION
S	15	SULFUR IN THIOETHERS AND MERCAPTANS	CB	37	CARBON AS IN BENZENE, PYRROLE	CL-	90	CHLORIDE ANION
S=C	16	TERMINAL SULFUR DOUBLY BONDED TO CARBON	NPYD	38	NITROGEN, AS IN PYRIDINE	BR-	91	BROMIDE ANION
S=O	17	SULFUR IN SULFOXIDES	NPYL	39	NITROGEN, AS IN PYRROLE	LI+	92	LITHIUM CATION
>S=N	17	SULFUR, TRICOORD, DOUBLY BONDED TO N	NC=C	40	NITROGEN ON N-C=C	NA+	93	SODIUM CATION
SO2	18	SULFUR IN SULFONES	NC=N	40	NITROGEN IN N-C=N	K+	94	POTASSIUM CATION
SO2N	18	SULFUR IN SULFONAMIDES	NC=P	40	NITROGEN IN N-C=P	ZINC	95	DIPOSITIVE ZINC
SO3	18	SULFONATE SULFUR	NC%C	40	NITROGEN ATTACHED TO C-C TRIPLE BOND	ZN+2	95	DIPOSITIVE ZINC
SO4	18	SULFATE SULFUR	CO2M	41	CARBOXYLATE ANION CARBON	CA+2	96	DIPOSITIVE CALCIUM
=SO2	18	SULFONE SULFUR DOUBLY BONDED TO CARBON	CS2M	41	CARBON IN THIOCARBOXYLATE ANION	CU+1	97	MONOPOSITIVE COPPER
SNO	18	SULFUR IN NITROGEN ANALOG OF A SULFONE	NSP	42	NITROGEN, TRIPLE BONDED	CU+2	98	DIPOSITIVE COPPER
SI	19	SILICON	NSO2	43	NITROGEN IN SULFONAMIDES	MG+2	99	DIPOSITIVE MAGNESIUM CATION
CR4R	20	CARBON IN 4-MEMBERED RINGS	NSO3	43	NITROGEN IN SULFONAMIDES, THREE O'S ON S			
HOR	21	HYDROGEN IN ALCOHOLS						

212 (!) different atom types,
grouped into 95 categories



```
MMFFMolProperties mmffMolProperties(ROMol &mol, std::string mmffVariant = "MMFF94",  
    boost::uint8_t verbosity = MMFF_VERBOSITY_NONE, std::ostream &oStream = std::cout);
```

```
// g++ -Wall -Wno-unused-function -o assignAtomTypes assignAtomTypes.cpp \  
// -I$RDBASE/Code -L$RDBASE/lib -lFileParsers -lRDGeneral -lForceFieldHelpers
```

```
#include <GraphMol/RDKitBase.h>  
#include <GraphMol/SmilesParse/SmilesParse.h>  
#include <GraphMol/ForceFieldHelpers/MMFF/AtomTyper.h>
```

```
using namespace RDKit;
```

```
int main()
```

```
{
```

```
    // assign and print MMFF atom types & charges
```

```
    ROMol *mol = SmilesToMol("CC(=O)C");
```

```
    MMFF::MMFFMolProperties mp(*mol);
```

```
    for (unsigned int i = 0; i < mol->getNumAtoms(); ++i) {
```

```
        std::cout << i + 1 << "\t" <<
```

```
            (unsigned int)mp.getMMFFAtomType(i) << "\t" <<
```

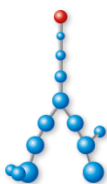
```
            mp.getMMFFFormalCharge(i) << "\t" <<
```

```
            mp.getMMFFPartialCharge(i) << std::endl;
```

```
    }
```

```
    return 0;
```

```
}
```

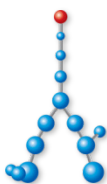


```
pyMMFFMolProperties = MMFFGetMoleculeProperties(mol, mmffVariant = "MMFF94", mmffVerbosity = 0)
```

```
# assign and print MMFF atom types
```

```
from rdkit import Chem
from rdkit.Chem import ChemicalForceFields

mol = Chem.MolFromSmiles("CC(=O)C")
mp = ChemicalForceFields.MMFFGetMoleculeProperties(mol)
for i in range(0, mol.GetNumAtoms()):
    print i + 1, '\t', mp.GetMMFFAtomType(i), \
          '\t', float(mp.GetMMFFFormalCharge(i)), \
          '\t', float(mp.GetMMFFPartialCharge(i))
```



Get MMFF atom types/charges

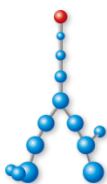
```
const boost::uint8_t mmffMolProperties.getMMFFAtomType(const unsigned int idx)  
const double mmffMolProperties.getMMFFFormalCharge(const unsigned int idx)  
const double mmffMolProperties.getMMFFPartialCharge(const unsigned int idx)
```

Include/exclude energy terms

```
void mmffMolProperties.setMMFFBondTerm(const bool state)  
void mmffMolProperties.setMMFFAngleTerm(const bool state)  
void mmffMolProperties.setMMFFStretchBendTerm(const bool state)  
void mmffMolProperties.setMMFFOopTerm(const bool state)  
void mmffMolProperties.setMMFFTorsionTerm(const bool state)  
void mmffMolProperties.setMMFFVdWTerm(const bool state)  
void mmffMolProperties.setMMFFEleTerm(const bool state)
```

Set MMFF94(s) variant, dielectric constant/model, verbosity

```
void mmffMolProperties.setMMFFVariant(const std::string mmffVariant)  
void mmffMolProperties.setMMFFDielectricConstant(const double dielConst)  
void mmffMolProperties.setMMFFDielectricModel(boost::uint8_t dielModel)  
void mmffMolProperties.setMMFFVerbosity(boost::uint8_t verbosity)  
void mmffMolProperties.setMMFFOStream(std::ostream *oStream)
```



Get MMFF atom types/charges

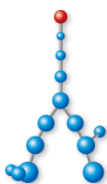
```
atomType = pyMMFFMolProperties.GetMMFFAtomType(idx)
formalCharge = pyMMFFMolProperties.GetMMFFFormalCharge(idx)
partialCharge = pyMMFFMolProperties.GetMMFFPartialCharge(idx)
```

Include/exclude energy terms

```
pyMMFFMolProperties.SetMMFFBondTerm(state)
pyMMFFMolProperties.SetMMFFAngleTerm(state)
pyMMFFMolProperties.SetMMFFStretchBendTerm(state)
pyMMFFMolProperties.SetMMFFOopTerm(state)
pyMMFFMolProperties.SetMMFFTorsionTerm(state)
pyMMFFMolProperties.SetMMFFVdWTerm(state)
pyMMFFMolProperties.SetMMFFEleTerm(state)
```

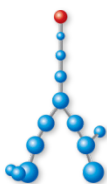
Set MMFF94(s) variant, dielectric constant/model, verbosity

```
pyMMFFMolProperties.SetMMFFVariant(mmffVariant)
pyMMFFMolProperties.SetMMFFDielectricConstant(dielConst)
pyMMFFMolProperties.SetMMFFDielectricModel(dielModel)
pyMMFFMolProperties.SetMMFFVerbosity(verbosity)
```



```
// g++ -Wall -Wno-unused-function -o computeEnergy computeEnergy.cpp \  
// -I$RDBASE/Code -L$RDBASE/lib -lFileParsers -lRDGeneral -lForceFieldHelpers -lForceField  
  
#include <GraphMol/RDKitBase.h>  
#include <GraphMol/FileParsers/MolSupplier.h>  
#include <GraphMol/FileParsers/MolWriters.h>  
#include <GraphMol/ForceFieldHelpers/MMFF/AtomTyper.h>  
#include <GraphMol/ForceFieldHelpers/MMFF/Builder.h>  
#include <ForceField/ForceField.h>  
  
using namespace RDKit;  
  
int main() {  
    std::string sdf = "ref_e2.sdf";  
    SDMolSupplier supplier(sdf, true, false);  
    SDWriter::SDWriter sdfWrite (sdf.substr(0, sdf.size() - 4) + "_MMFF_min.sdf");  
    ROMol *mol;  
    std::cout << "MMFF ENERGY\tBEFORE MIN\tAFTER MIN\n";  
    for (unsigned int i = 0; i < supplier.length(); ++i) {  
        mol = supplier[i];  
        std::string molName = "";  
        if (mol->hasProp("_Name")) {  
            mol->getProp("_Name", molName);  
        }  
        MMFF::MMFFMolProperties mp(*mol);  
        ForceFields::ForceField *field = MMFF::constructForceField(*mol, &mp);  
        field->initialize();  
        double e1 = field->calcEnergy();  
        field->minimize();  
        double e2 = field->calcEnergy();  
        std::cout << molName << std::setprecision(3) << std::fixed << "\t\t" << e1 << "\t\t" << e2 << "\n";  
        sdfWrite.write(*mol);  
        delete mol;  
    }  
    sdfWrite.close();  
    return 0;  
}
```

Load dataset from SDF;
minimize;
compute MMFF energy
(C++)



```
from rdkit import Chem
from rdkit.Chem import AllChem

sdf = 'ref_e2.sdf'
supplier = Chem.SDMolSupplier(sdf, True, False)
sdfWrite = Chem.SDWriter(sdf[:-4] + '_MMFF_min.sdf')

print 'MMFF ENERGY\tBEFORE MIN\tAFTER MIN'
for i in range(0, len(supplier)):
    mol = supplier[i]
    if (mol.HasProp('_Name')):
        molName = mol.GetProp('_Name')
        mp = AllChem.MMFFGetMoleculeProperties(mol)
        field = AllChem.MMFFGetMoleculeForceField(mol, mp)
        e1 = field.CalcEnergy()
        field.Minimize()
        e2 = field.CalcEnergy()
        print '{0:}\t\t{1:.3f}\t\t{2:.3f}'.format(molName, e1, e2)
    sdfWrite.write(mol)
sdfWrite.close()
```

Load dataset from SDF;
minimize;
compute MMFF energy
(Python)

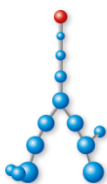
```
// g++ -Wall -Wno-unused-function -o gen3D gen3D.cpp -I$RDBASE/Code -L$RDBASE/lib \  
// -lFileParsers -lRDGeneral -lForceFieldHelpers -lForceField -lDistGeomHelpers
```

```
#include <GraphMol/RDKitBase.h>  
#include <GraphMol/FileParsers/MolSupplier.h>  
#include <GraphMol/FileParsers/MolWriters.h>  
#include <GraphMol/ForceFieldHelpers/MMFF/AtomTyper.h>  
#include <GraphMol/ForceFieldHelpers/MMFF/Builder.h>  
#include <GraphMol/MolOps.h>  
#include <GraphMol/DistGeomHelpers/Embedder.h>  
#include <ForceField/ForceField.h>
```

```
using namespace RDKit;
```

```
int main() {  
    std::string smi = "ref_e2.smi";  
    SmilesMolSupplier supplier(smi);  
    SDWriter::SDWriter sdfWrite (smi.substr(0, smi.size() - 4) + "_MMFF_gen3D.sdf");  
    ROMol *mol, *molH;  
    std::cout << "NAME\t\tMMFF ENERGY\n";  
    for (unsigned int i = 0; i < supplier.length(); ++i) {  
        mol = supplier[i];  
        std::string molName = "";  
        if (mol->hasProp("_Name")) {  
            mol->getProp("_Name", molName);  
        }  
        molH = MolOps::addHs(*mol);  
        DGeomHelpers::EmbedMolecule(*molH);  
        MMFF::MMFFMolProperties mp(*molH);  
        ForceFields::ForceField *field = MMFF::constructForceField(*molH, &mp);  
        field->initialize();  
        field->minimize();  
        double e = field->calcEnergy();  
        std::cout << molName << std::setprecision(3) << std::fixed << "\t\t" << e << "\n";  
        sdfWrite.write(*molH);  
        delete mol;  
        delete molH;  
    }  
    sdfWrite.close();  
    return 0;  
}
```

Load dataset from SMILES;
generate 3D;
MMFF-minimize
(C++)



```
from rdkit import Chem
from rdkit.Chem import AllChem

smi = 'ref_e2.smi'
supplier = Chem.SmilesMolSupplier(smi)
sdfWrite = Chem.SDWriter(smi[:-4] + '_MMFF_gen3D.sdf')

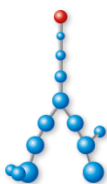
print 'NAME\t\tMMFF ENERGY'
for i in range(0, len(supplier)):
    mol = supplier[i]
    if (mol.HasProp('_Name')):
        molName = mol.GetProp('_Name')
        molH = AllChem.AddHs(mol)
        AllChem.EmbedMolecule(molH)
        mp = AllChem.MMFFGetMoleculeProperties(molH)
        field = AllChem.MMFFGetMoleculeForceField(molH, mp)
        field.Minimize()
        e = field.CalcEnergy()
        print '{0:}\t\t{1:.3f}'.format(molName, e)
    sdfWrite.write(molH)
sdfWrite.close()
```

Load dataset from SMILES;
generate 3D;
MMFF-minimize
(Python)

The RDKit implementation was validated against the official **MMFF94** (761 molecules) and **MMFF94s** (265 molecules) validation suites available in the CCL data archives (both dative and hypervalent notations):

<http://ccl.net/chemistry/resources/data/index.shtml>

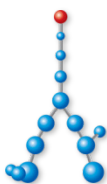
- All atoms are assigned correct MMFF types and charges
- All force-field terms are assigned correct force constants
- All MMFF energies are correctly computed
- Gradients were also validated against the **TINKER** MMFF implementation
- Even better, atom type, charge and force constants are correctly assigned also starting from SMILES representations of the validation suite molecules
- MMFF validation is included as part of the RDKit test suite
(Code/ForceField/MMFF/testMMFFForceField.cpp)



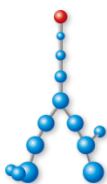
- GBSA implicit solvation model

During energy minimization, support for:

- Fixed atoms
- Harmonic constraints on selected atoms/groups
- Internal coordinate constraints
(in addition to distances, also angles and torsions)




- The MMFF implementation
- UFF gains something, too
- MMFF applications



(Lack of) Planarity of some generated atomic coordinates after minimisation.

 greglandrum is assigned

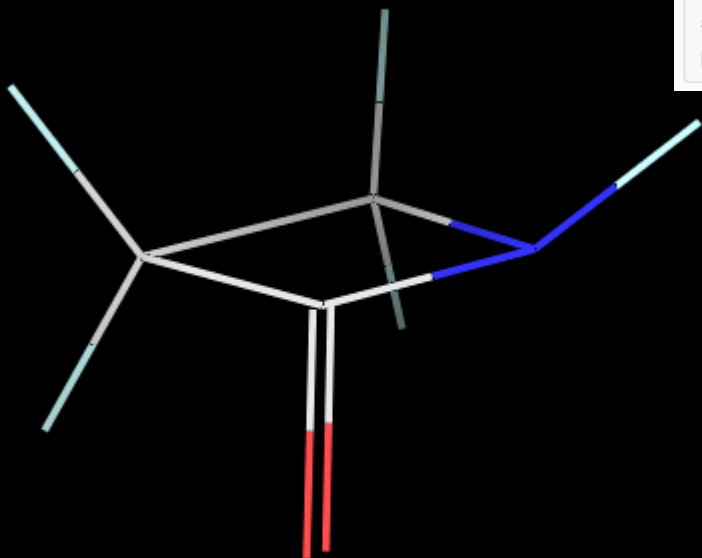
Milestone: 2013_09_1 

The problem is the lack of an inversion term in the RDKit implementation of UFF. This is definitely solvable (UFF has an inversion term, we just never implemented it), but it will probably take some time.

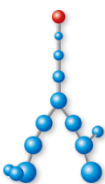
```
#!/usr/bin/env python

from rdkit import Chem
from rdkit.Chem import AllChem

# a simple mol in a simple world
m = Chem.MolFromSmiles("C1C(=O)NC1")
# the usual recipe
Chem.AddHs(m)
AllChem.EmbedMolecule(m)
# Opt
AllChem.UFFOptimizeMolecule(m)
# Double bond O should be planar - it is not
print >>file('wrong_structure.sdf', 'w'), Chem.MolToMolBlock(m)
```

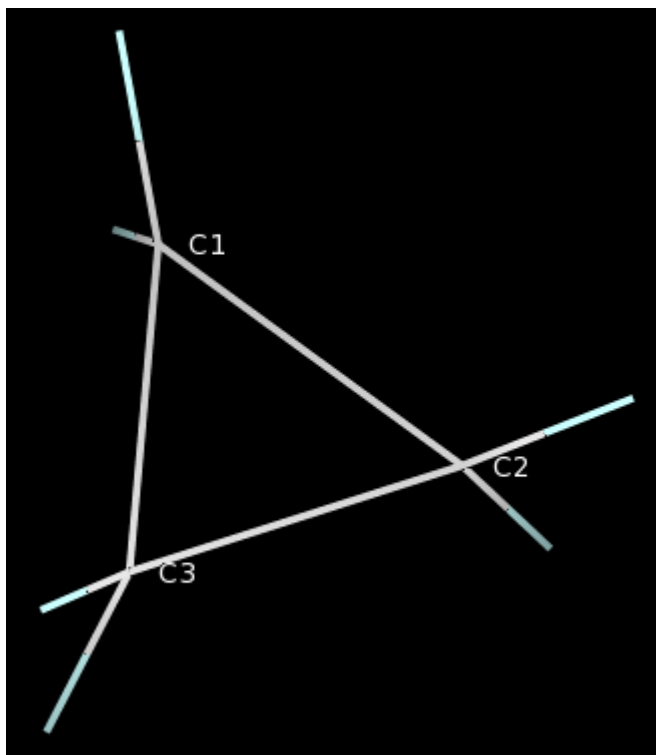


- The issue is arguably connected with the lack of the inversion term
- Back-porting the OOP term from MMFF to UFF should most likely fix things

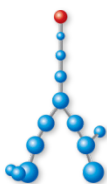


```
PyMOL>get_angle id 1, id 2, id 3  
cmd.get_angle: 56.674 degrees.  
PyMOL>get_angle id 1, id 3, id 2  
cmd.get_angle: 66.654 degrees.  
PyMOL>get_angle id 2, id 1, id 3  
cmd.get_angle: 56.672 degrees.
```

```
PyMOL>get_distance id 1, id 2  
cmd.get_distance: 1.610 Angstroms.  
PyMOL>get_distance id 1, id 3  
cmd.get_distance: 1.465 Angstroms.  
PyMOL>get_distance id 2, id 3  
cmd.get_distance: 1.465 Angstroms.
```

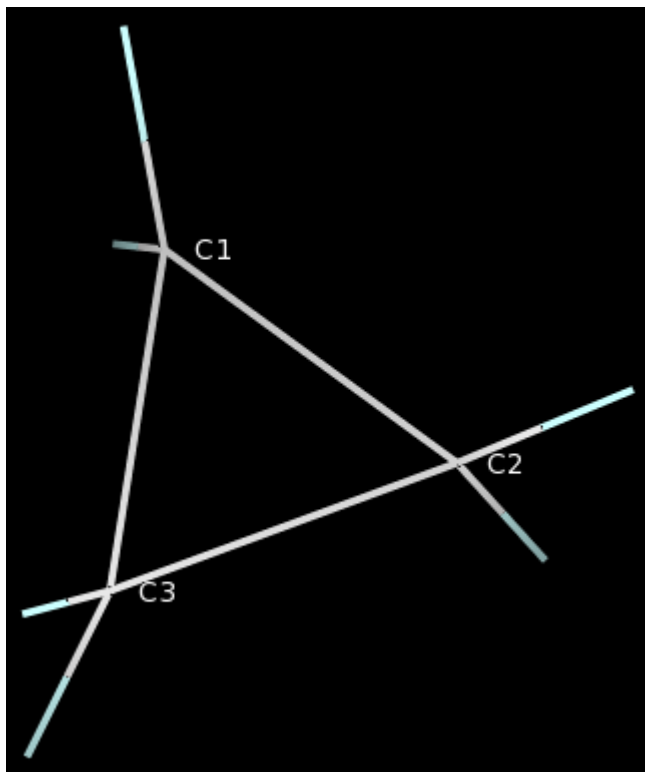


- I noticed that also geometries of small rings containing only sp^3 atoms were somehow ill
- Angle enumeration in UFF was implemented using the same `neighborMatrix` used for non-bonded interactions
- However, since atoms **C1**, **C3** are connected both directly and through atom **C2** and the matrix has only one node per atom pair, some angle terms may be missing



```
PyMOL>get_angle id 1, id 2, id 3  
cmd.get_angle: 60.003 degrees.  
PyMOL>get_angle id 1, id 3, id 2  
cmd.get_angle: 59.998 degrees.  
PyMOL>get_angle id 2, id 1, id 3  
cmd.get_angle: 59.999 degrees.
```

```
PyMOL>get_distance id 1, id 2  
cmd.get_distance: 1.517 Angstroms.  
PyMOL>get_distance id 1, id 3  
cmd.get_distance: 1.517 Angstroms.  
PyMOL>get_distance id 2, id 3  
cmd.get_distance: 1.517 Angstroms.
```

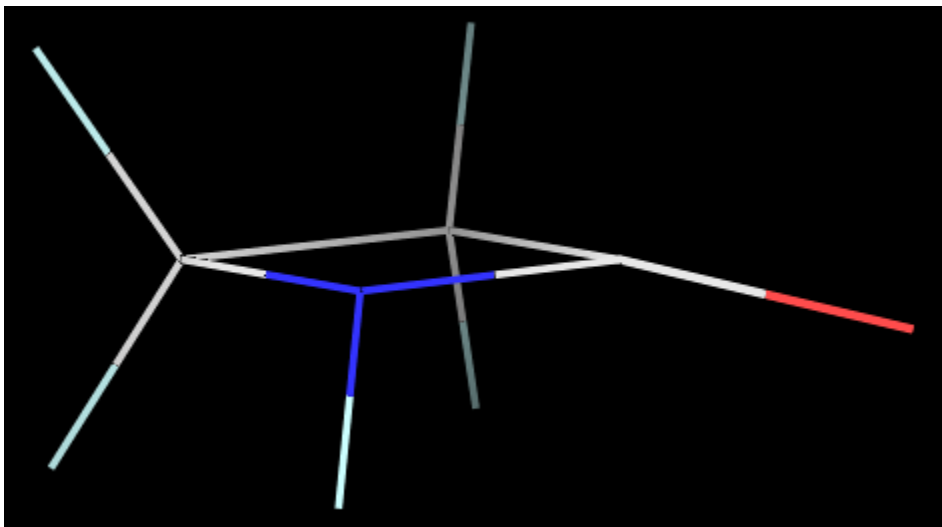


- Both in MMFF and UFF a graph-based method is now used for angle enumeration, while `neighborMatrix` is used only for non-bonded interactions
- This also allows reducing the size of `neighborMatrix` nodes from a 32-bit int to 2 bits, resulting in a 256-fold lower RAM usage
- All torsions are now included also in 3-membered rings
- Angles and distances now look OK

- Then, I implemented inversions according to the original paper by Rappé et al.:

UFF, a Full Periodic Table Force Field for Molecular Mechanics and Molecular Dynamics Simulations

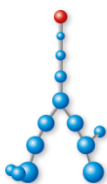
A. K. Rappé,* C. J. Casewit,[†] K. S. Colwell, W. A. Goddard III,[#] and W. M. Skiff[†]



- After making sure that the inversion code was working properly, I ran the script which triggered issue 62:

```
# a simple mol in a simple world
m = Chem.MolFromSmiles("C1C(=O)NC1")
m2 = Chem.AddHs(m)
AllChem.EmbedMolecule(m2)
pyFF = AllChem.UFFGetMoleculeForceField(m2)
pyFF.Minimize()
print >>file('uff_beta-lactam.sdf','w'), \
    Chem.MolToMolBlock(m2)
```

The geometry improved slightly,
but was still far from being satisfactory! Very disappointing.



- Investigations on the details of the UFF implementation revealed minor discrepancies with respect to Rappé's paper, which anyway had no impact on the geometries of small rings containing sp^2 atoms

Typos and comments for UFF

There are some obvious typos in the UFF paper, and I believe there are a few subtle ones as well. Here I list places where my implementation does not completely agree with what is written in the UFF paper.

- Equation (2) of [Rappe et al. 1992](#) is written as follows.

$$r_{IJ} = r_I + r_J + r_{BO} + r_{EN}$$

However, this method does not result in agreement with their published equilibrium bond lengths. Anthony Rappe informed me that this equation is in error and I have instead implemented the following (beginning with Version 4.4.2).

$$r_{IJ} = r_I + r_J + r_{BO} - r_{EN}$$

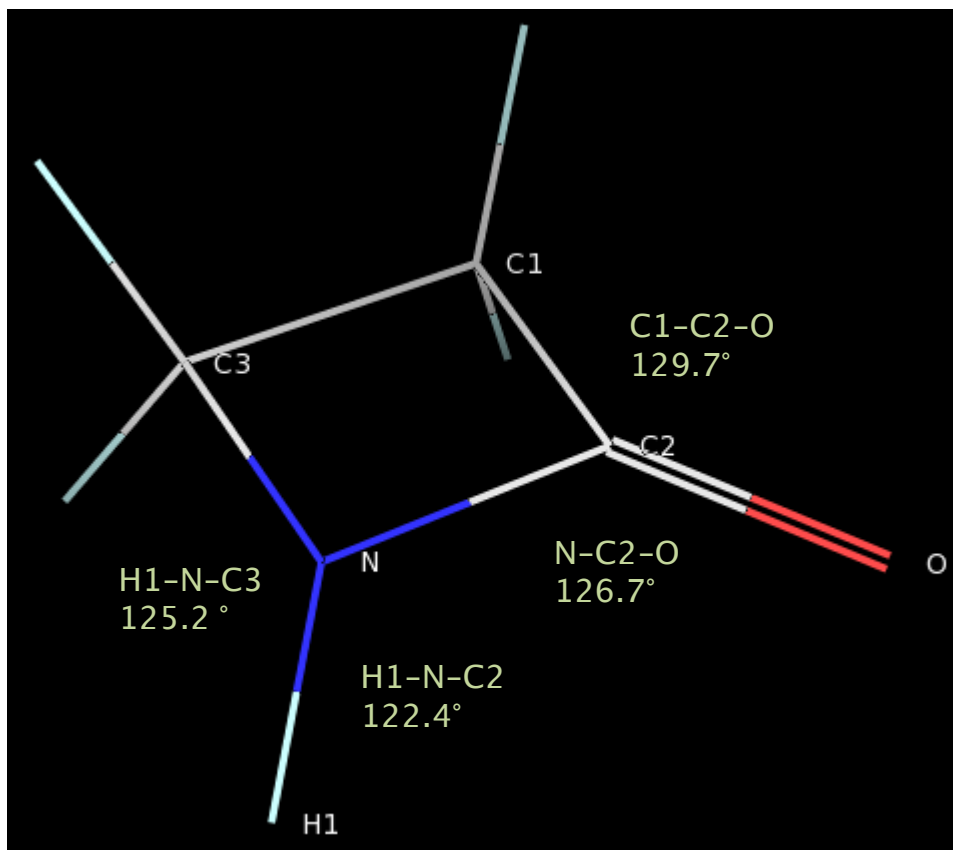
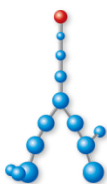
- Equation (13) of [Rappe et al. 1992](#) is written with some mistakes in the superscripts and subscripts. Here is the equation as implemented into Towhee (beginning with Version 4.4.2).

$$K_{IJK} = \text{beta} (Z_I^* Z_K^* / r_{IK}^5) r_{IJ} r_{JK} [3 r_{IJ} r_{JK} (1 - \text{Cos}^2(\text{theta}_0)) - r_{IK}^2 \text{Cos}(\text{theta}_0)]$$

- Equation 10 and the preceding text in [Rappe et al. 1992](#) does not accurately reflect the implementation of bending angles in UFF. For the linear case the equation should actually read

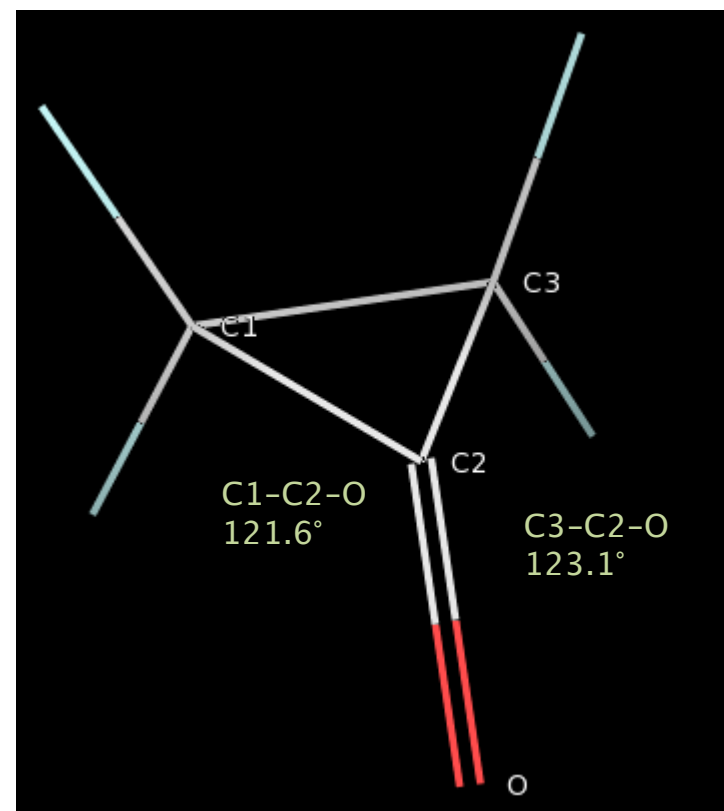
$$U = K_{IJK}/n^2 [1 + \text{Cos}(n \text{ theta})]$$

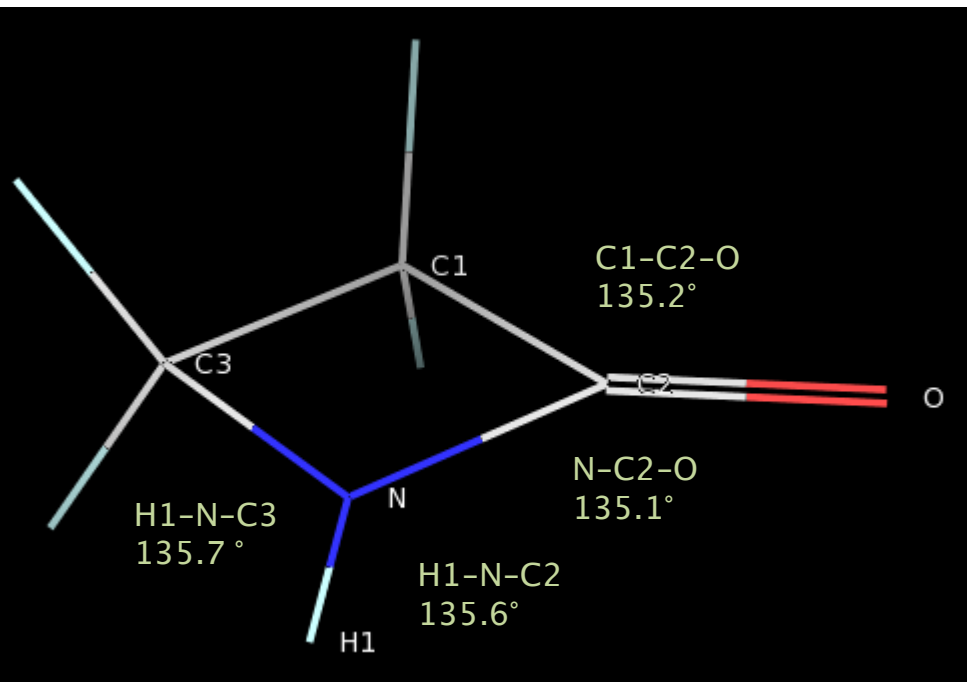
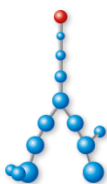
- During my investigations I also ran across these suggestions by Marcus Martin, the author of [Towhee](#), who found some typos/mistakes in the original paper. However, none of these fixes solved issue 62



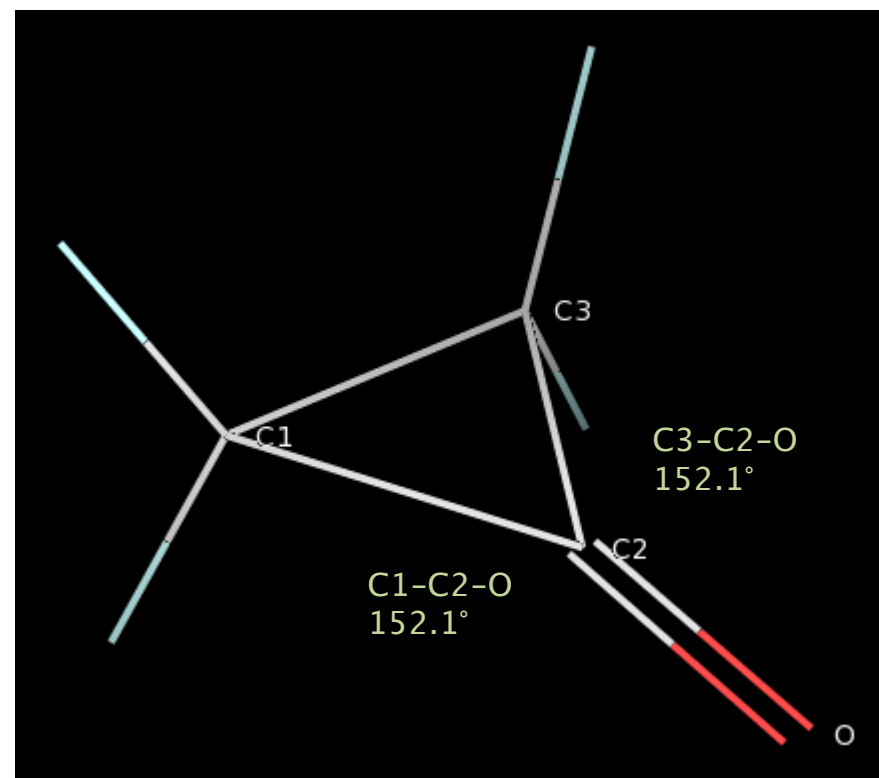
- There seems to be a **competition** between angle and OOP bend terms, the first pulling the exocyclic atom out-of-plane, the second trying to keep it coplanar with the ring

- Measuring the angles involving two edges of the small ring and an exocyclic atom shows that such angles are close to **120°**, which is the equilibrium angle for **sp²** atoms





- I set equilibrium values for **exocyclic** angles involving **sp²** atoms to **135°** and **150°**, and **endocyclic** angles to **90°** and **60°** for 4- and 3-membered rings, respectively



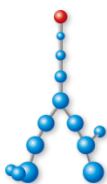
- Now exocyclic **sp²** atoms are coplanar with the ring, as they should be
- Please note that this hack impacts only on **sp²** atoms in 3- and 4-membered rings

Small ring geometries obtained with original UFF, hacked UFF, AM1

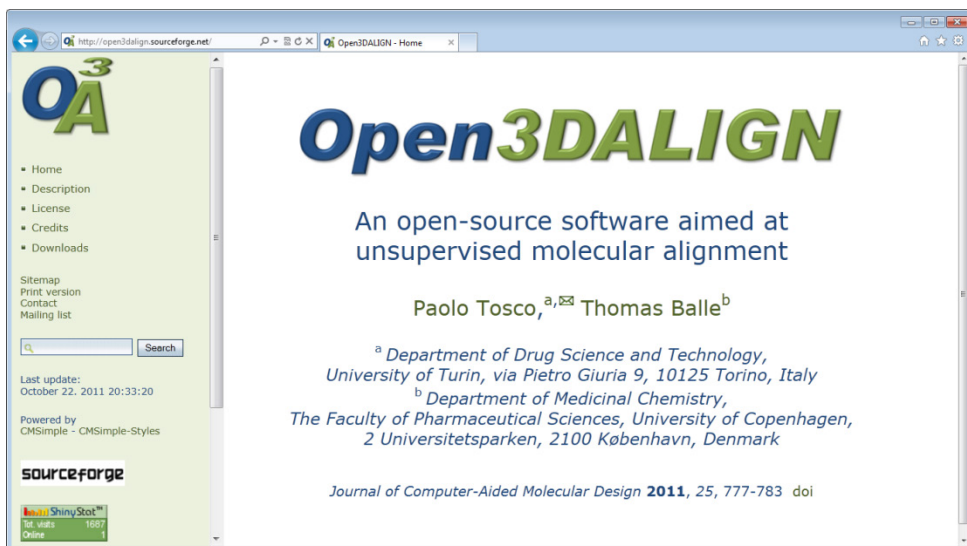
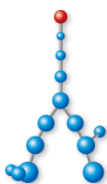
	original UFF	hacked UFF	AM1
1			
2			
3			
4			
5			

	original UFF	hacked UFF	AM1
6			
7			
8			
9			
10			

	original UFF	hacked UFF	AM1
11			
12			
13			
14			

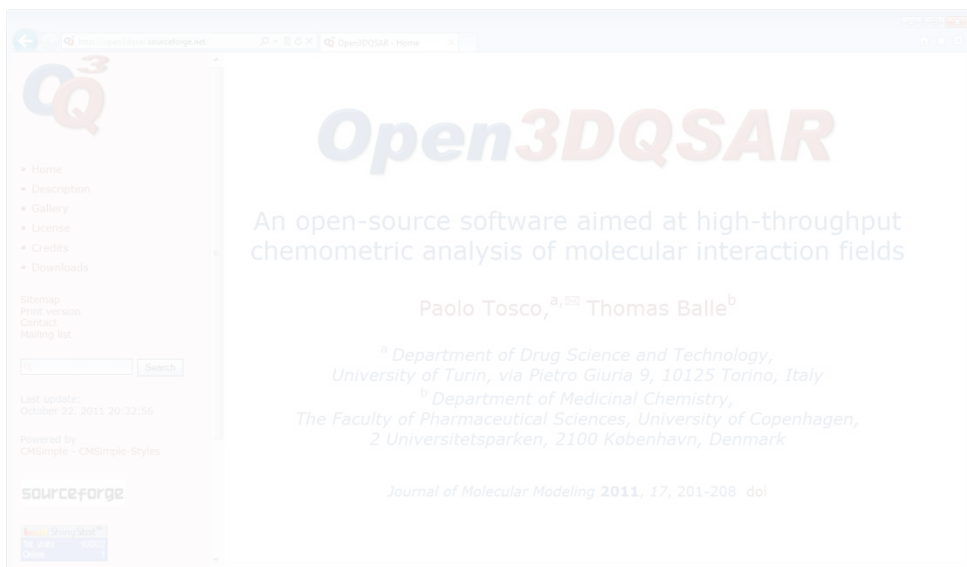


- The MMFF implementation
- UFF gains something, too
- **MMFF applications**



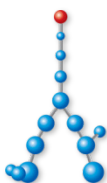
• Open3DALIGN

- molecular alignment
 - single/multiple conformations
 - atom-based alignment
 - single-run



• Open3DQSAR

- MIF computation
- PLS model building and validation
- variable selection



Open3DALIGN uses a 2-step alignment strategy

1

- a cost function c_{ij} is computed for pairing heavy atoms i, j between reference (R) and probe (P) molecules

	c_{00}	c_{01}	c_{02}	c_{03}	c_{04}	c_{05}	c_{06}	c_{07}	c_{08}	c_{09}
	c_{10}	c_{11}	c_{12}	c_{13}	c_{14}	c_{15}	c_{16}	c_{17}	c_{18}	c_{19}
i	c_{20}	c_{21}	c_{22}	c_{23}	c_{24}	c_{25}	c_{26}	c_{27}	c_{28}	c_{29}
	c_{30}	c_{31}	c_{32}	c_{33}	c_{34}	c_{35}	c_{36}	c_{37}	c_{38}	c_{39}
	c_{40}	c_{41}	c_{42}	c_{43}	c_{44}	c_{45}	c_{46}	c_{47}	c_{48}	c_{49}
	c_{50}	c_{51}	c_{52}	c_{53}	c_{54}	c_{55}	c_{56}	c_{57}	c_{58}	c_{59}
	j									

$$c_{ij} = w_1 |q(R)_i - q(P)_j| + w_2 s_{ij} + \sum_{k=1}^K \frac{(h(P)_{ik} - h(R)_{jk})^2}{h(P)_{ik} + h(R)_{jk}}$$

$q(R)_i, q(P)_j$

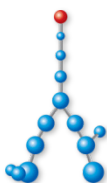
MMFF94 charges of atoms

s_{ij}

degree of chemical similarity between MMFF94 atom types i and j

$h(R), h(P)$

K -binned histograms whose k -th bin represents the number of atoms in R and P lying in a $(k-1, k)$ distance range from the i -th and j -th atom, respectively



Open3DALIGN uses a 2-step alignment strategy

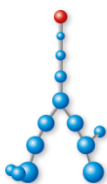
1

- The total matching cost is minimized via the Jonker-Volgenant algorithm yielding an **E** matrix which describes intermolecular atom pairings

<i>i</i>	c₀₀	c₀₁	c₀₂	c₀₃	c₀₄	c₀₅	c₀₆	c₀₇	c₀₈	c₀₉
c₁₀	c ₁₀	c ₁₁	c ₁₂	c ₁₃	c ₁₄	c ₁₅	c ₁₆	c ₁₇	c ₁₈	c ₁₉
c₂₀	c ₂₀	c ₂₁	c ₂₂	c ₂₃	c ₂₄	c ₂₅	c ₂₆	c ₂₇	c ₂₈	c ₂₉
c₃₀	c ₃₀	c ₃₁	c ₃₂	c ₃₃	c ₃₄	c ₃₅	c ₃₆	c ₃₇	c ₃₈	c ₃₉
c₄₀	c ₄₀	c ₄₁	c ₄₂	c ₄₃	c ₄₄	c ₄₅	c ₄₆	c ₄₇	c ₄₈	c ₄₉
c₅₀	c ₅₀	c ₅₁	c ₅₂	c ₅₃	c ₅₄	c ₅₅	c ₅₆	c ₅₇	c ₅₈	c ₅₉
	<i>j</i>									

- The **E** matrix is then ranked and trimmed to give a **D** matrix, whose atom pairs are least-square-fitted

<i>i</i>	<i>j</i>		<i>i</i>	<i>j</i>
4	0		5	7
2	2		1	8
0	3		3	5
3	5		2	2
5	7			
1	8			



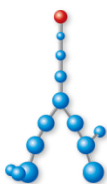
Open3DALIGN uses a 2-step alignment strategy

2

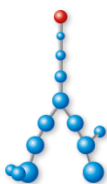
- This initial, coarse alignment is then iteratively refined via the **SDM** algorithm and scored
- Optionally, cost function parameters may be iteratively optimized until the best scoring alignment is found

Scoring function:

$$s = \sum_{k=1}^p \left(\alpha + \frac{1 + \beta \cdot |q(R)_{D[k,0]} + q(P)_{D[k,1]}|}{1 + |q(R)_{D[k,0]} - q(P)_{D[k,1]}|} \right) \cdot \exp(-\gamma \cdot r_{D[k,0]D[k,1]}^2)$$

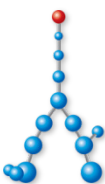


- The RDKit already implements least-square-fit and 3D transformation functions dedicated to molecular alignment:
- ```
double getAlignmentTransform (const ROMol &prbMol, const ROMol &refMol, RDGeom::Transform3D
&trans, int prbCid = -1, int refCid = -1, const MatchVectType *atomMap = 0, const
RDNumeric::DoubleVector *weights = 0, bool reflect = false, unsigned int maxIters = 50)
```
- ```
double alignMol (ROMol &prbMol, const ROMol &refMol, int prbCid = -1, int refCid = -1, const  
MatchVectType *atomMap = 0, const RDNumeric::DoubleVector *weights = 0, bool reflect = false,  
unsigned int maxIters = 50)
```
- All we need is to generate via the **Open3DALIGN** functionality a MatchVectType with matching atom pairs and a RDNumeric::DoubleVector with weights, and pass them to the least-square-fit function



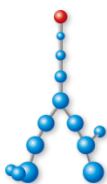
```
O3A o3a(ROMol &prbMol, const ROMol &refMol, MMFF::MMFFMolProperties *prbMP,  
        MMFF::MMFFMolProperties *refMP, const int prbCid = -1, const int refCid = -1,  
        const bool reflect = false, const unsigned int maxIters = 50,  
        const unsigned int options = O3_USE_MMFF_WEIGHTS, LAP *extLAP = NULL)
```

```
// align probe onto reference (returns RMSD)  
double rmsd = o3a.align()  
  
// get RMSD and transformation which overlays probe and reference molecules  
std::pair<double, RDGeom::Transform3D *> rmsdTrans = o3a.trans()  
  
// get the O3AScore value of the alignment (no need to actually align coordinates)  
double o3aScore = o3a.score()  
  
// get the MatchVectType vector  
const RDKit::MatchVectType *matches = o3a.matches()  
  
// get the weight vector  
const RDNumeric::DoubleVector *weights = o3a.weights()
```



```
GetO3A(prbMol, refMol, prbPyMMFFMolProperties, refPyMMFFMolProperties,  
       prbCid = -1, refCid = -1, reflect = False, maxIters = 50, options = O3_USE_MMFF_WEIGHTS)
```

```
# align probe onto reference (returns RMSD)  
rmsd = o3a.Align()  
  
# get RMSD and transformation which overlays probe and reference molecules  
(rmsd, trans) = o3a.Trans()  
  
# get the O3AScore value of the alignment (no need to actually align coordinates)  
o3aScore = o3a.Score()  
  
# get the MatchVectType vector  
matches = o3a.Matches()  
  
# get the weight vector  
weights = o3a.Weights()
```



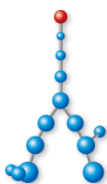
```
// g++ -Wall -Wno-unused-function -o o3aAlign o3aAlign.cpp -I$RDBASE/Code \  
// -L$RDBASE/lib -lFileParsers -lRDGeneral -lForceFieldHelpers -lForceField -lMolAlign
```

```
#include <GraphMol/RDKitBase.h>  
#include <GraphMol/FileParsers/MolSupplier.h>  
#include <GraphMol/FileParsers/MolWriters.h>  
#include <GraphMol/ForceFieldHelpers/MMFF/AtomTyper.h>  
#include "GraphMol/MolAlign/AlignMolecules.h"  
#include "GraphMol/MolAlign/O3AAlignMolecules.h"
```

```
using namespace RDKit;
```

```
int main() {  
    std::string sdf = "ref_e2_scrambled.sdf";  
    std::string o3aSdf = "ref_e2_O3A.sdf";  
    SDMolSupplier supplier(sdf, true, false);  
    int nMol = supplier.length();  
    const int refNum = 48;  
    ROMol refMol = *(supplier[refNum]);  
    MMFF::MMFFMolProperties refMP(refMol);  
    SDWriter::SDWriter *o3aMol = new SDWriter::SDWriter(o3aSdf);  
    std::cout << "N\t\tSCORE\t\tRMSD" << std::endl;  
    for (int prbNum = 0; prbNum < nMol; ++prbNum) {  
        ROMol prbMol = *(supplier[prbNum]);  
        MMFF::MMFFMolProperties prbMP(prbMol);  
        MolAlign::O3A o3a(prbMol, refMol, &prbMP, &refMP);  
        std::cout << prbNum + 1 << "\t\t" << std::fixed << std::setprecision(2)  
            << o3a.score() << "\t\t" << o3a.align() << std::endl;  
        o3aMol->write(prbMol);  
    }  
    o3aMol->close();  
    return 0;  
}
```

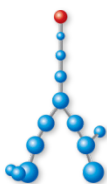
Align all molecules in a SDF file
on a common reference (C++)



```
from rdkit import Chem
from rdkit.Chem import AllChem

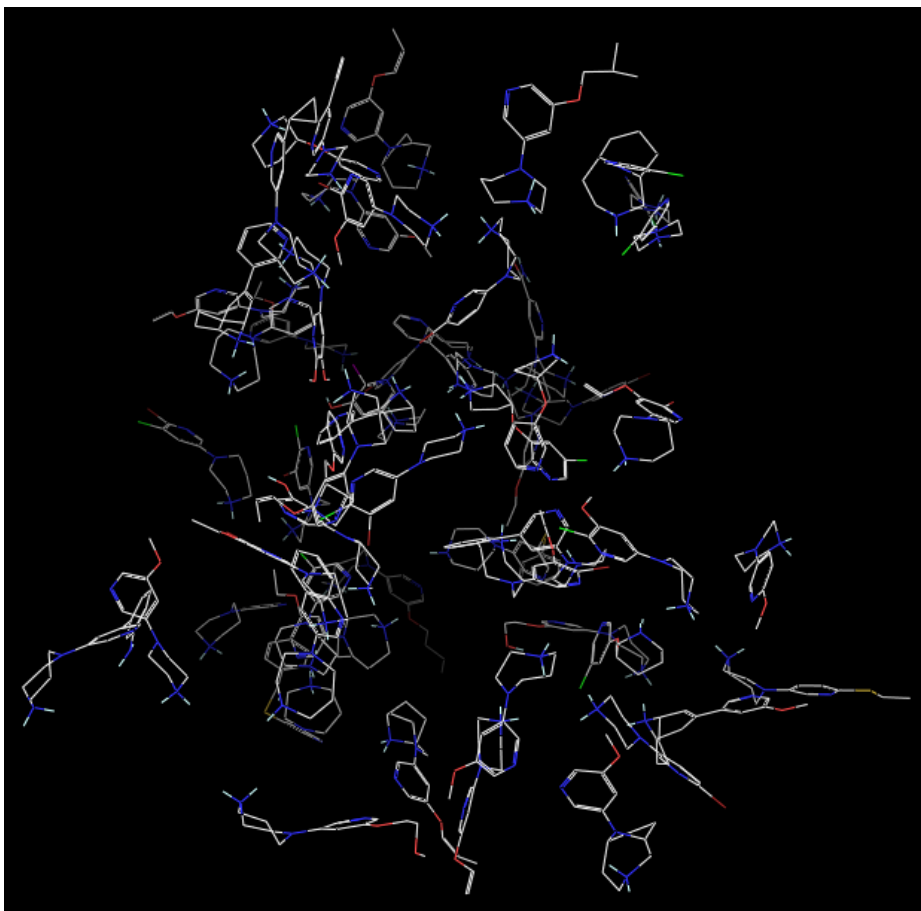
sdf = 'ref_e2.sdf'
o3aSdf = 'ref_e2_PyO3A.sdf'
supplier = Chem.SDMolSupplier(sdf, True, False)
nMol = len(supplier)
refNum = 48
refMol = supplier[refNum]
refPyMP = AllChem.MMFFGetMoleculeProperties(refMol)
o3aMol = Chem.SDWriter(o3aSdf)
print 'N\t\tSCORE\t\tRMSD'
for prbNum in range(0, nMol):
    prbMol = supplier[prbNum]
    prbPyMP = AllChem.MMFFGetMoleculeProperties(prbMol)
    pyO3A = AllChem.GetO3A \
        (prbMol, refMol, prbPyMP, refPyMP)
    print '{0:}\t\t{1:.2f}\t\t{2:.2f}'.format(prbNum + 1, pyO3A.Score(), pyO3A.Align())
    o3aMol.write(prbMol)
o3aMol.close()
```

Align all molecules in a SDF file
on a common reference (Python)

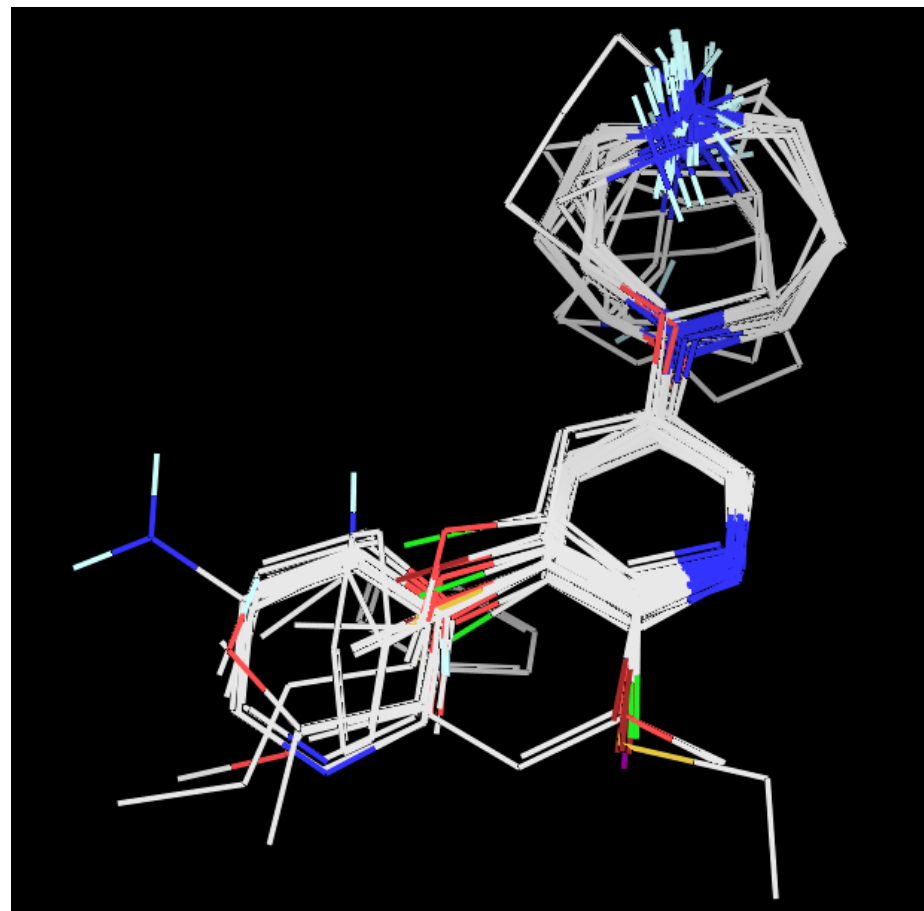


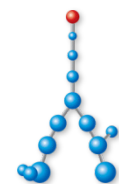
O3A alignment: before and after

Before

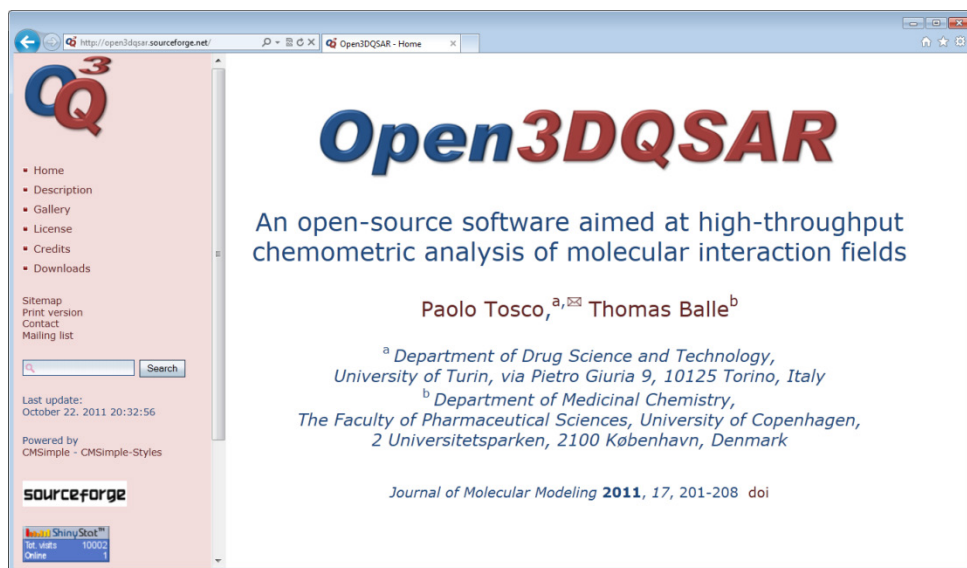


After

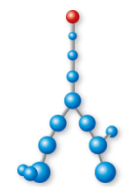




- Methods to perform OOTB multi-conformational and multi-threaded alignment
- Iterative refinement of overlays
- Implementation of **O3Q** functionality



- **Open3DQSAR**
 - MIF computation
 - PLS model building and validation
 - variable selection



Thanks

- To Nik and Greg
- To NIBR for funding
- To all scientists who publish their science in such an open, clear and truly reproducible form as Halgren did with MMFF